

Original Article

Elucidation of the RNA-granule inducing sodium azide stress response through transcriptome analysis

Mani Garg, Gopalakrishna Poornima, Purusharth I. Rajyaguru*

Department of Biochemistry, Indian Institute of Science, Bangalore 560012, India



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ABSTRACT

Sodium azide is a commonly used cytochrome oxidase inhibitor that leads to translation repression and RNA granule assembly. The global changes in mRNA abundance in response to this stressor are unknown. RGG-motif proteins Scd6 and Sbp1 are translation-repressors and decapping-activators that localize to and affect the assembly of RNA granules in response to sodium azide stress. Transcriptome-wide effects of these proteins remain unknown. To address this, we have sequenced transcriptome of the: a) wild type strain under unstressed and sodium azide stress, b) $\Delta scd6$ and $\Delta sbp1$ strains under unstressed and sodium azide stress. Transcriptome analysis identified altered abundance of many transcripts belonging to stress-responsive pathways which were further validated by qRT-PCR results. Abundance of several transcripts was altered in $\Delta scd6/\Delta sbp1$ under normal conditions and upon stress. Overall, this study provides critical insights into transcriptome changes in response to sodium azide stress and the role of RGG-motif proteins in these changes.

1. Introduction

The cellular response to stress is a feature of all kingdoms of life. The ability of cells to sense and respond to several kinds of environmental stress is essential for survival and evolutionary progress. The ultimate aim of the stress response is to alter the cellular proteome to efficiently combat the stress. This involves upregulating stress-responsive genes and downregulating various energy-consuming pathways, including mRNA translation. Rewiring mRNA abundance, which in turn affects the proteome, is critical for cellular stress response pathways [1–3].

The specific genes and signaling pathways altered in response to stress may vary depending on the type of stressor. A thorough study of the differential network of genes affected by various stressors is key to understand the molecular mechanisms underlying the stress response machinery. Oxidative stress has been implicated in the development and progression of many diseases such as cancer, diabetes, and cardiovascular and neurodegenerative disorders [4–7]. Sodium azide is a well-known oxidative stress-inducing agent that inhibits cytochrome oxidase to impair the mitochondrial electron transport chain and induces oxidative stress [8,9]. Sodium azide has been used to study various biological processes such as the fermentative ability of yeast grown under different oxygen conditions [10]. Interestingly, it has also been used as a weak mutagen in yeast protoplasts [11]. It increases

thermo-tolerance of glucose-grown yeast [12] and is routinely used in studies of mRNA translation control since sodium azide treatment leads to general translation repression [13]. Despite the knowledge that sodium azide can inhibit protein and RNA synthesis [14,15], the global impact of sodium azide on mRNA abundance has not been characterized.

Global mRNA translation repression has emerged as an important hallmark of stress responses. The resulting non-translating mRNAs assemble into conserved membraneless mRNA-protein (mRNP) compartments known as RNA granules. Stress granules and P-bodies are well-known examples of RNA granules implicated in translation repression [16,17]. Along with non-translating mRNAs, stress granules and P-bodies are also sites of mRNA storage and decay [16,18]. RNA granule-dynamics are greatly influenced by a number of different categories of proteins, some of which include RNA-binding proteins, translation repressors, and mRNA decay proteins. RNA granules can assemble in response to a wide array of stresses such as glucose deprivation, amino acid starvation, oxidative stress (sodium azide), heat shock, and stationary phase [19–22].

A subset of RGG-motif proteins have been identified as translation repressors that target eIF4G1, a conserved translation initiation factor [23]. Scd6 and Sbp1 are two such proteins with multiple RG-/RGG-repeats. Scd6 is a conserved RGG-motif protein with orthologs identified in most model systems, including humans [24]. Along with the

* Corresponding author.

E-mail address: rajyaguru@iisc.ac.in (P.I. Rajyaguru).

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RGG-motif, these two proteins have different RNA binding domains. Scd6 contains an N-terminal Lsm domain, a central FDF-domain, and a C-terminal RGG-motif [24]. The RGG-motif of Sbp1 is flanked by two RRM-domains, one on either side [25]. Both the proteins can affect mRNA stability by activating decapping [26,27]. Deletion of Scd6 and Edc3 (Enhancer Of Decapping 3) leads to a synthetic growth defect [28]. *MFA2* mRNA decapping was defective in the absence of both Edc3 and Scd6, suggesting that the two proteins participate in decapping activation. Consistent with this idea, Scd6 co-localizes with Edc3 and Dcp2 in RNA granules such as processing bodies (P-bodies), which are the sites of mRNA decay [23,29]. Sbp1, on the other hand, was identified as a multicopy suppressor of decapping defects in a conditional mutant of Dcp1 and Dcp2 [27]. Sbp1, like Scd6, localizes to P-bodies and perturbation of Sbp1 levels perturbs P-body assembly [27]. Consistent with the presence of RNA-binding domains, mRNAs associate with Scd6 and Sbp1 in vivo [30,31]. As decapping activators, Scd6 and Sbp1 affect transcript abundance by altering mRNA stability, although only *MFA2* has been identified as an mRNA target so far. Given that sodium azide treatment induces P-body formation and localization of Scd6 and Sbp1 to P-bodies [23,25], it is likely to modulate the role of Scd6 and Sbp1 in maintaining transcript stability. However, the effect of Scd6 and Sbp1 on the transcriptome under normal and sodium azide treated conditions has not been assessed.

mRNAs bound by the translation repression complex are recruited to mRNP granules and are protected from decay machinery. Consequently, they can be stored for a long period of time through a process known as ‘masking’, an evolutionarily conserved mechanism. For example in worms and flies, certain key mRNAs encoding developmental proteins are stored and protected from translation and decay [18]. *Tral*, the *Drosophila* ortholog of Scd6, is involved in masking and storage of nanos mRNA and regulates the Gurken secretory pathway in conjunction with Me31B (Dhh1 in yeast) and Cup. Further, *Tral* mutants show dysregulated embryonic development [32]. In worms, CAR-1 (Scd6 ortholog), along with CGH-1 (Dhh1 ortholog), is crucial for germline granule maintenance [33]. As such, the deletion of Scd6 and Sbp1 could render certain sequestered mRNAs unprotected, thereby promoting their decay. Investigating mRNA abundance in Δ *scd6* and Δ *sbp1* strains can provide insight about specific mRNAs downregulated in response to the loss of masking by translation repression. Such response is likely to be altered upon exposure to stress, for example sodium azide.

Keeping the above in focus, this work aimed to a) understand the impact of sodium azide on the yeast transcriptome and b) identify the transcripts affected by the RGG-motif proteins Scd6 and Sbp1 in response to sodium azide stress. Our results show that sodium azide treatment leads to widespread changes in transcript abundance in *S. cerevisiae* wild type strain (BY4741). Many functional categories of transcripts with altered expression overlap with those reported in response to other oxidative stresses. We further identified multiple transcripts whose abundance was altered in normal growth conditions in the absence of Scd6 and Sbp1 and in response to sodium azide stress. It is possible that some transcripts identified in this study are direct mRNA targets of Scd6 and Sbp1. In summary, this work provides a detailed insight into the gene expression changes induced by sodium azide stress and also sheds light on the role of Scd6/Sbp1 in maintaining transcript abundance under normal conditions and in response to sodium azide stress.

2. Materials and methods

2.1. Growth conditions

Yeast strains used in this study are listed in Supplementary Table 1. Strains were grown at 30°C in yeast extract-peptone (YP) medium supplemented with 2% glucose. For secondary culture, cells were diluted ($OD_{600} \sim 0.1$) in YP + 2% glucose. At mid-log phase (OD_{600} 0.4–0.5),

cultures were divided into two; one was treated with 0.5% sodium azide (NaN_3), and the other with water (control) for 30 min. A 20% NaN_3 stock solution was made in water and added to culture to obtain a final concentration of 0.5% NaN_3 . Post-treatment, cells were harvested by centrifugation and used for further analysis immediately or stored at $-80^\circ C$ for further experiments.

2.2. Microscopy

The PAB1GFP strain was grown in YP + 2% glucose at 30 °C (until $OD_{600} \sim 0.4$ – 0.5), followed by 0.5% NaN_3 treatment for 30 min. Cells were centrifuged at 14,000 rpm for 12 s and pellets were resuspended in 20 μ l of supernatant media. A total of 5 μ l of the cell suspension was spotted on coverslips for live cell imaging. All images were acquired using a Deltavision Elite microscope system with softWoRx 3.5.1 software (Applied Precision, LLC) and an Olympus 100 \times , oil-immersion 1.4 NA objective. Exposure time and transmittance settings for the Green Fluorescent Protein (GFP) channel were 0.2 s and 32%, respectively. Images were captured as 512 \times 512-pixel files with a CoolSnapHQ camera (Photometrics) using 1 \times 1 binning for yeast. All the images were deconvolved using standard softWoRx deconvolution algorithms. ImageJ was used to adjust all images to equalize contrast ranges.

2.3. RNA isolation and analysis

RNA was isolated from two biological replicates for each test sample using the hot acidic phenol protocol described in Current Protocols in Molecular Biology (1996) Unit 13.12 [34]. Briefly, the cell pellet was washed with nuclease free water once and resuspended thoroughly in TES (10 mM TrisCl pH 7.5, 10 mM EDTA and 0.5% SDS). 400 μ l acidic phenol pH 4.5 (0.1 M citrate buffer saturated) was added to the suspension and incubated at 65 °C for 60 min with occasional vortexing. Samples were spun at 14,000 rpm for 10 min at 4 °C. The top aqueous layer was transferred to a fresh tube. Subsequently, 400 μ l chloroform was added, mixed, and spun as above. The aqueous layer was taken and mixed with 1/10th volume 3 M NaAc pH 5.2 and 2.5 volumes of 100% ethanol. The samples were precipitated at $-80^\circ C$ overnight, then spun as above and the pellet was washed with 70% ethanol, dried, and resuspended in 50 μ l nuclease free water. RNA quality was checked by 1.2% agarose formamide gel electrophoresis. RNA quality and quantity were also assessed using the Agilent Bioanalyzer RNA 6000 Nano kit (Agilent Technologies) and the Qubit HS RNA kit (Thermo Fisher Scientific). Samples were sent to Clevergene for RNA-seq.

2.4. RNA library preparation and sequencing

One μ g of total RNA was enriched for mRNA with the NEBNext Poly (A) mRNA magnetic isolation Module (E7490, New England Biolabs, USA) according to the manufacturer's instructions. The protocol uses polyT magnetic beads to enrich mRNA molecules with polyA tails. The polyA enriched mRNA was used for library preparation. The NEBNext Ultra II RNA Library Prep Kit for Illumina (E7770, New England Biolabs, USA) was used for library preparation following the manufacturer's protocol. In brief, enriched mRNA was fragmented and converted to cDNA followed by second-strand synthesis. NEBNext sample purification beads were used to isolate double-stranded cDNA. After end repair, hairpin adapters were ligated to the purified cDNA and the hairpin structure was cleaved using USER enzyme. NEBNext Multiplex Oligos for Illumina (E7335S, New England Biolabs, USA) were used for polymerase chain reaction (PCR) amplification to generate indexed sequencing libraries. The PCR conditions were 98 °C for 1 min for initial denaturation, followed by 7 cycles of 98 °C for 10 s, 65 °C for 75 s, and a final extension at 65 °C for 5 min. The amplicons were purified using NEBNext sample purification beads.

Library quality and quantity was checked using an Agilent

Bioanalyzer 2100 with DNA 1000 kits and a Qubit dsDNA HS Assay Kit (Thermo Fisher Scientific), and libraries were sequenced on an Illumina HiSeqX to generate 150 bp paired-end reads.

2.5. Transcriptome analysis

The QC passed reads were mapped to the indexed *Saccharomyces cerevisiae* reference genome (S288c) using STAR v2 aligner [35]. PCR and optical duplicates were marked and removed using Picard tools. Gene expression level values were obtained as read counts using the featureCounts program [36]. Differential expression analysis was carried out using the edgeR package [37] after normalizing the data based on the trimmed mean of M (TMM) values. After normalization, 248 features (3.76%) were removed from the analysis because they did not have at least 1 counts-per-million in at least two samples.

Enrichment analysis for Biological process, Molecular function, Cellular component and KEGG Pathway analysis was performed using the ClusterProfiler R package [38]. Gene Ontology (GO) and pathway terms with adjusted p -value $\leq .05$ were considered significant. The GPlot R package was used to visualize GO enrichment results [39]. KEGG pathway visualization was performed by the pathView R Bioconductor package [40]. The submitted raw data can be accessed here (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE145162>).

2.6. Reverse transcription and quantitative real-time PCR

For target validation, isolated RNA was first subjected to DNaseI treatment (Thermo, EN0525) by combining 8 μ g of total RNA, 4 units of DNaseI, and DNaseI Buffer with MgCl₂ in a 30 μ l reaction. After incubating for 30 min 37 °C, DNaseI was inactivated with 3 μ l of 50 mM EDTA for 10 mins at 65 °C. Following DNaseI treatment, RNA quality was checked again using 1.2% agarose formamide gel electrophoresis. DNaseI treated RNA was then used for cDNA synthesis.

One microgram of RNA was used to synthesize cDNA using the RevertAid RT Reverse Transcription Kit (Thermo, K1691) according to the manufacturer's protocol. cDNA was diluted 1:10 and real-time PCR was performed using TB Green™ Premix Ex Taq™ (TaKaRa). For qRT-PCR, three technical replicates were assembled with 2 μ l cDNA/reaction and 0.5 μ M each primer in a BioRad iQ5 Real-Time PCR Detection System. Primers used for qRT-PCR are listed in Supplementary Table 2. The PCR conditions were 95 °C for 12 min for initial denaturation, followed by 30 cycles of 95 °C for 20 s, 46 °C for 30 s, and 72 °C for 30 s. DNA was quantified in every cycle at the extension step. Melt curve acquisition was carried out at 64 °C for 8 s. Ct values were extracted with auto baseline and manual threshold. $\Delta\Delta$ Ct method was used to calculate the final log₂ FoldChange values, which were then plotted on a box and whisker plot using GraphPad prism 7.0. Significance was calculated by a ratio paired t -test.

3. Results

3.1. RNA sequencing analysis of wild type, Δ scd6, and Δ sbp1 strains

Wild type, Δ scd6, and Δ sbp1 strains were grown in duplicates (biological replicates) under normal conditions overnight. Secondary cultures were inoculated at 0.1 OD₆₀₀ and allowed to grow for two generations. Cultures were then divided in half and one sample was treated with 0.5% sodium azide and the other with water, followed by incubation at 30 °C for 30 min (Fig. 1a). To ensure that sodium azide stress was induced under these conditions, a strain expressing Pab1 (a conserved stress granule marker) endogenously tagged with GFP was treated with sodium azide in an identical manner and observed under a fluorescence microscope to visualize stress granule formation. Pab1-GFP foci were induced in cells treated with sodium azide but not in those treated with an equivalent volume of water, confirming that the stress granules were indeed induced by sodium azide treatment

(Fig. 1b).

RNA was isolated from treated and untreated wild type, Δ scd6, and Δ sbp1 strains and subjected to denaturing formamide agarose gel electrophoresis (Fig. 1c). PolyA enriched mRNA was used for cDNA synthesis and library preparation. Illumina sequencing was performed on each sample with two biological replicates. Sequencing data quality was checked using FastQC and MultiQC software. The data was checked for base call quality distribution, % bases above Q20, Q30, % GC, and sequencing adapter contamination, and all samples passed the QC threshold (Q30 > 90%). On average, 90.69% of the QC-passed reads aligned to the reference genome. After removing PCR and optical duplicates, gene expression values were obtained as read counts using featureCounts software. Expression similarity between biological replicates was checked by Spearman correlation and the co-efficient value between replicates was > 0.97.

For differential expression analysis, biological replicates were grouped as “Control” and “Test” with the control category listed first in all comparisons (for example: Wild type Unt vs NaN₃ Treated considers Wild type Unt as control and NaN₃ Treated as test). Genes with an absolute log₂ fold change ≥ 2 and p -value $\leq .05$ were considered significantly differentially expressed (Fig. 2). A summary of differentially expressed genes is presented as volcano plots and heatmaps (Fig. 3 and Supplementary Figs. 1 & 2).

3.2. Stress-response genes show significant changes in transcript abundance in response to sodium azide treatment

RNA-seq analysis revealed that 270 genes were differentially expressed following sodium azide treatment, of which 200 were upregulated and 70 were downregulated (Fig. 2). Genes belonging to the carbohydrate metabolic process gene ontology (GO) category constituted the largest (22 genes) category of upregulated genes in sodium azide treated wild type cells (Fig. 4a). Two other GO categories upregulated upon sodium azide stress include oxidative stress-responsive genes (11 genes) and abiotic stress-responsive genes (15 genes). This is consistent with the action of sodium azide on cytochrome oxidase and on cellular redox balance. The downregulated genes were prominently represented in the GO category of ribosome biogenesis (17 genes) and cell division (13 genes) (Fig. 4a). The downregulation of ribosome biogenesis genes was consistent with previously reported observations that sodium azide treatment globally represses translation.

Interestingly, two non-coding RNAs HRA1 (Hidden-in-Reading frame Antisense-1) and RUF21 (RNA of Unknown Function 21) are downregulated in wild type cells following sodium azide treatment (Supplementary Fig. 3). Interestingly these RNAs were downregulated in Δ sbp1 (Supplementary Fig. 3) cells also following sodium azide treatment. HRA1 is a substrate of RNase P and is speculated to be involved in 18S rRNA maturation [41].

3.3. Transcriptome changes in Δ scd6 and Δ sbp1 strains

In Δ scd6 background as compared to wild type, 53 genes were differentially expressed as compared to wild type under normal conditions; 28 genes were upregulated, and 25 genes were downregulated (Fig. 2). Of the upregulated genes, sugar transporter genes (including hexose transporter HXT7) were prominently represented (Fig. 4b). The downregulated genes belong to different GO categories including intron homing, heme-binding, and alpha-glucosidase activity. Three non-coding RNAs, RUF23, RUF5-1, and RUF5-2, were upregulated upon deletion of SCD6, indicating that Scd6 may play a role in maintaining non-coding RNA levels apart from mRNA levels (Supplementary Fig. 3).

RNA-seq analysis of the Δ sbp1 strain as compared to wild type revealed an altered abundance of 22 different transcripts. This included 10 upregulated and 12 downregulated genes (Fig. 2). Interestingly, the sugar transporter gene HXT7, which was upregulated in the absence of Scd6, was also upregulated upon Sbp1 deletion, indicating that both

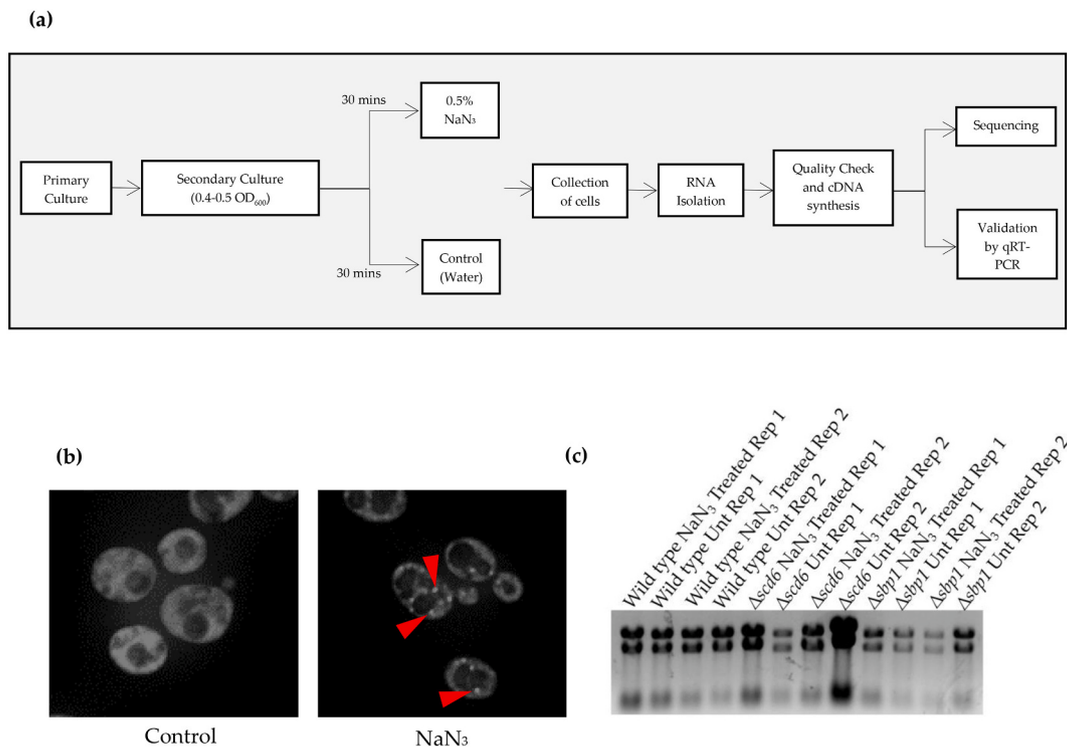


Fig. 1. (a) Schematic depicting the experimental workflow. Primary yeast cultures were grown for two biological replicates of every strain. For secondary culture, cells were diluted to 0.1 OD₆₀₀. At mid-log (0.4–0.5 OD₆₀₀), cells were treated with 0.5% NaN₃ for 30 min followed by collection of cells by centrifugation. For control cells, equivalent volume of water was added. RNA was isolated using hot phenol followed by quality check. Sequencing was carried out following cDNA synthesis. Differentially expressed genes were validated by qRT-PCR analysis which was carried out for three technical replicates of each biological replicates; (b) Live cell imaging of cells expressing genomically-tagged PAB1GFP strain treated with NaN₃ at the same time as the cells used for isolating the RNA used for RNA-sequencing. Arrows indicate the presence of stress granules; (c) Agarose formamide gel electrophoresis image of the RNA sent for RNA sequencing analysis. Rep 1 and Rep 2 refer to Replicate 1 and Replicate 2, respectively.

proteins could play a cooperative role in regulating the abundance of HXT7 mRNA. The downregulated genes include GO categories such as superoxide dismutase and copper ion binding (Fig. 4b).

Following sodium azide stress, in *Δscd6* strain as compared to wild type, 18 genes were differentially expressed with 9 genes being upregulated and an equal number of genes being downregulated (Fig. 2). GO categories such as chitin deacetylase activity and spermine transmembrane transporter activity were represented in the downregulated genes (Supplementary Fig. 4). Interestingly, GO categories including intron homing and cell division were represented in the upregulated genes (Supplementary Fig. 4). Following sodium azide stress, in *Δsbp1* strain as compared to wild type, there was differential expression of 64 genes, with 27 genes being upregulated and 37 genes being downregulated (Fig. 2). It is noteworthy that the intron homing genes (*All1*, *Al5_beta*, and *Scel1*) upregulated in the absence of *Scd6* were also upregulated in the absence of *Sbp1* upon oxidative stress, although to different extents (Supplementary Fig. 4). This indicated that both the RGG-motif proteins acted on similar mRNAs in this important biological process.

In response to sodium azide treatment in *Δscd6* and *Δsbp1* backgrounds as compared to untreated, abundance of 245 and 533 transcripts changed respectively (Fig. 2). Genes encoding proteins involved in carbohydrate metabolic process, oxidative and abiotic stress response were significantly enriched in the sodium azide treated *Δscd6* and *Δsbp1* cells (Supplementary Fig. 4b & c). Downregulation of cellular component genes of cell surface family is a common feature of sodium azide stress response mounted by wild type, *Δscd6* and *Δsbp1* strains (Fig. 4a, Supplementary Fig. 4b & c). Specifically, different aspartyl proteases like *MKC7* and *BAR1* were significantly reduced in response to sodium azide. In contrast to this, ribosome biogenesis and cell division related genes did not show any significant change in case of *Δscd6* background upon sodium azide stress as compared to untreated cells. In *Δscd6*

background, genes belonging to cell wall category were downregulated whereas in *Δsbp1* background genes belonging to helicase activity category were downregulated (Supplementary Fig. 4b). This is in contrast to the wild type response to sodium azide stress.

3.4. Validation of RNA-seq results by qRT-PCR

To choose a suitable control gene for validating the RNA-seq results, we checked the mRNA levels of three different control genes, namely *ACT1*, *SCR1*, and *PGK1*, under different conditions by qRT-PCR. In absence of stress, both *ACT1* and *SCR1* genes were not modulated significantly in *Δscd6* and *Δsbp1* strains (Supplementary Fig. 5). *SCR1* was used as a control to validate transcript levels in *Δscd6* and *Δsbp1* backgrounds under untreated condition; however, in response to sodium azide treatment, *SCR1* expression was significantly upregulated when compared with *ACT1* and *PGK1* (Supplementary Fig. 5). Therefore, *PGK1* was chosen as the control for mRNA target validation specific to sodium azide treatment.

We validated the change in abundance of transcripts that were differentially expressed upon sodium azide treatment in wild type yeast cells. We focused on *HSP30*, *SP11*, *TPO2*, *CIN5*, and *FDH1*. All the tested transcripts were upregulated in wild type cells upon sodium azide treatment (Fig. 5a). A comparison of the changes in transcript abundance measured by RNA-seq and qRT-PCR are presented (Supplementary Fig. 6a).

We performed qRT-PCR to validate the RNA-seq results for transcripts differentially expressed in *Δscd6*. *HXT7* and *CWP1* transcript levels were validated using *SCR1* as the internal control. The abundance of these transcripts increased in *Δscd6* (Fig. 5b and Supplementary Fig. 6b). Interestingly, both transcripts also increased in abundance in *Δsbp1* (Supplementary Fig. 6b & c). Based on these results, we conclude

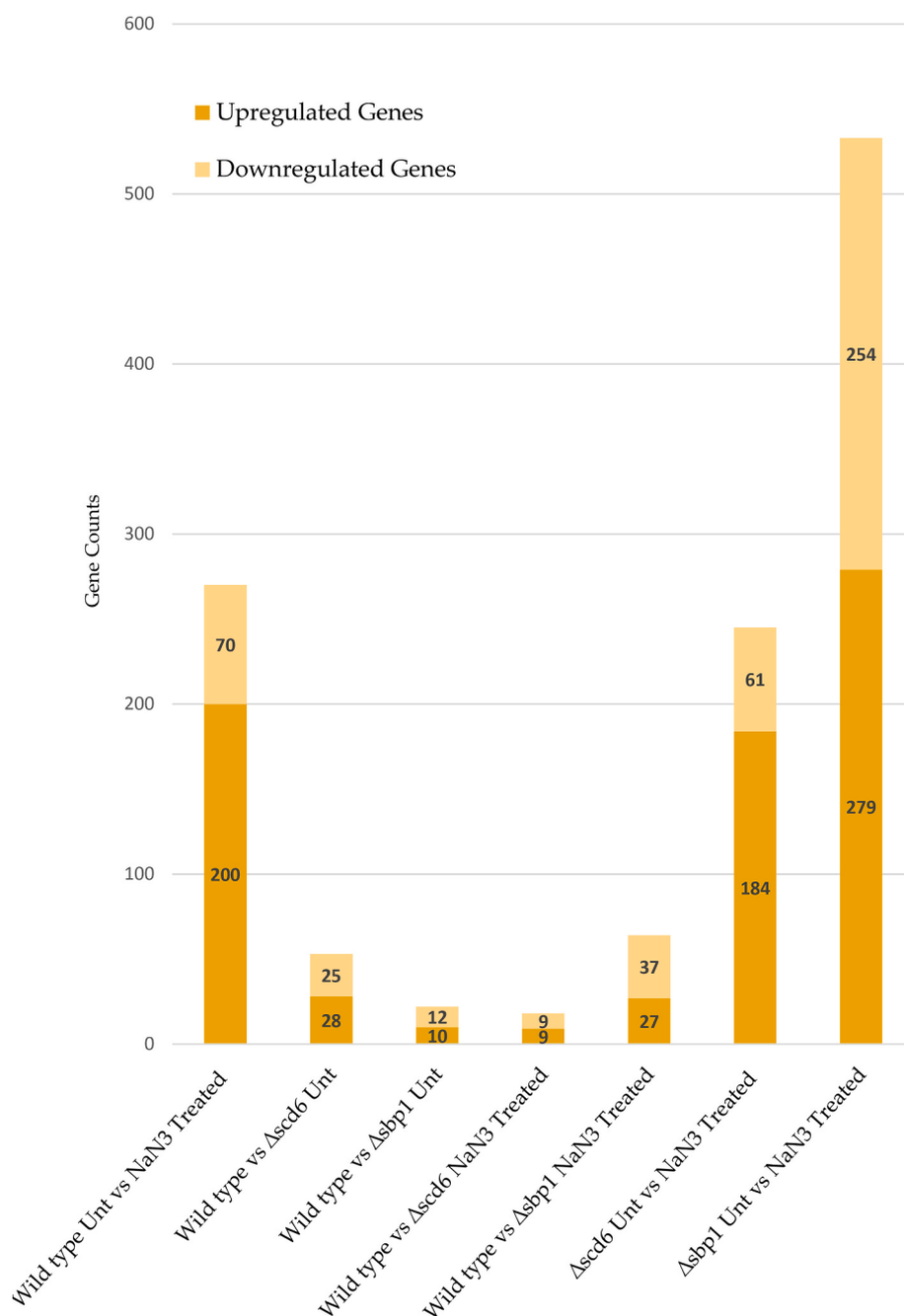


Fig. 2. Summary of significantly up and downregulated genes in different conditions.

that *Scd6* and *Sbp1* reduce the abundance of *HXT7* and *CWP1* transcripts.

Similarly, to validate RNAseq results for differentially expressed mRNAs in $\Delta sbp1$, candidate mRNAs such as *HXK1*, *GLK1*, *GSY1*, and *GPH1* were chosen. Two points motivated us to investigate these transcripts. First, RNA-seq results indicated that the levels of these mRNAs were significantly modulated. Second, *Sbp1* binds to these transcripts in yeast cells [30], indicating that these changes in transcript level are likely mediated by *Sbp1* binding to these mRNAs. The results of qRT-PCR showed increased abundance of these mRNAs (Fig. 5c and Supplementary Fig. 6c). Based on these results, we conclude that the deletion of *Sbp1* increases the abundance of *HXK1*, *GLK1*, *GSY1*, and *GPH1* transcripts, all of which encode enzymes involved in glucose metabolism.

Changes in transcript levels in $\Delta scd6$ and $\Delta sbp1$ cells upon sodium azide treatment as compared to wild type treated cells were also

validated. *HXK1*, *GPH1*, and *GSY1* were tested as all three transcripts showed increased abundance in RNA-seq data (Supplementary Fig. 7). We observed that all the tested transcripts showed increased levels by qRT-PCR in the absence of *Scd6* and *Sbp1* following sodium azide treatment, thus confirming the RNA-seq results. Taken together, these observations validate the RNA-seq results by qRT-PCR.

4. Discussion

Sodium azide represses global translation and induces RNA granule formation within minutes after cells are exposed to the stressor [15]. This in turn, could affect the half lives of many transcripts, thereby contributing to altered gene expression profile. However, the subset of genes affected by sodium azide stress has remained unknown. The present study used RNA sequencing to analyze the change in mRNA levels of yeast cells exposed to sodium azide stress. Our study also

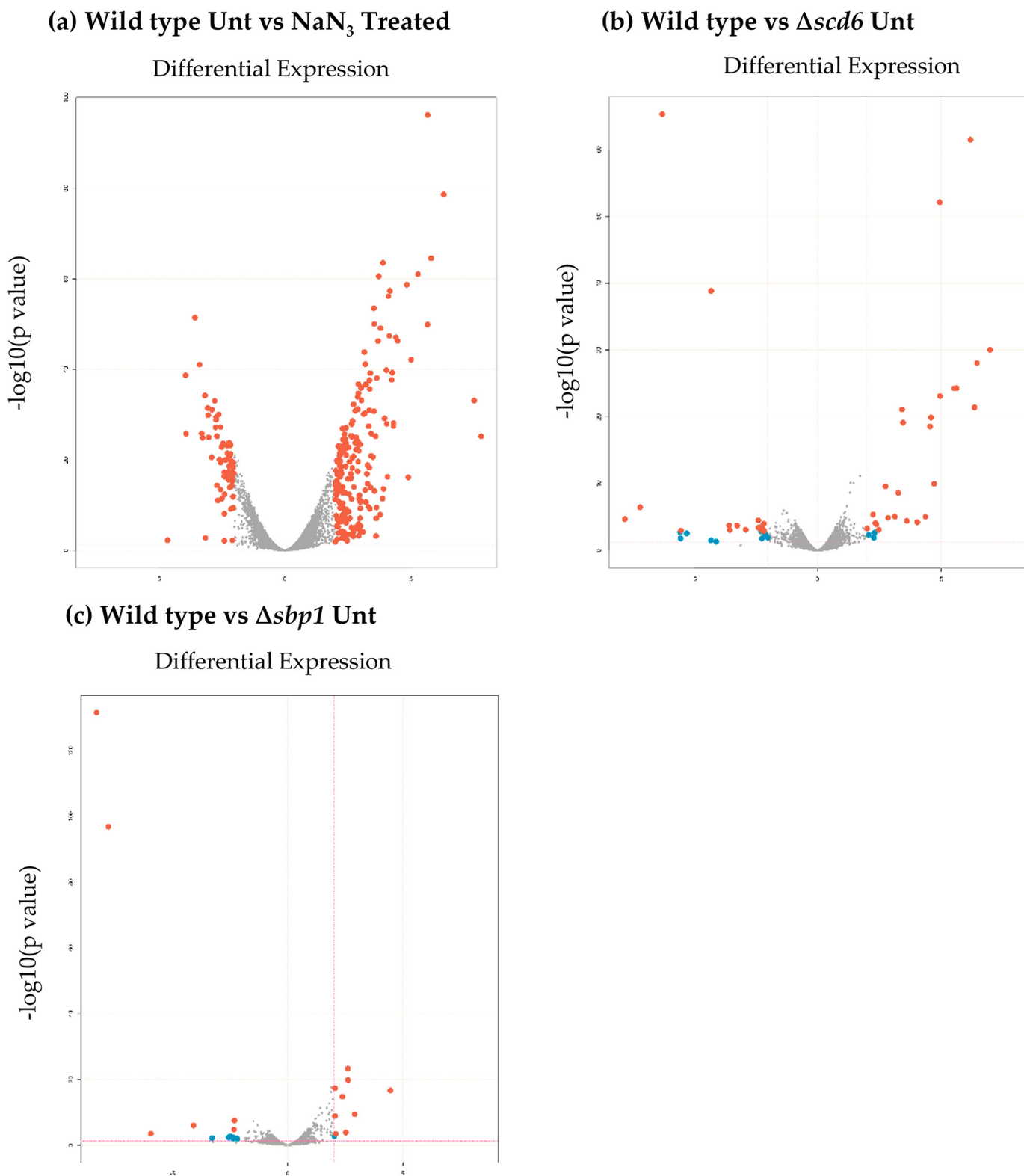


Fig. 3. Volcano plot showing changes in transcript levels in: (a) wild type cells upon sodium azide treatment, (b) wild type cells upon deletion of *Scd6* and, (c) wild type cells upon deletion of *Sbp1*. Red dots indicate absolute log₂fold change ≥ 2 and FDR/adjusted *p*-value ≤ .05. Blue dots indicate log₂fold change ≥ 2 and *p*-value ≤ .05. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

evaluated the mRNA abundance changes upon deletion of RGG-motif containing decapping activators such as *Scd6* and *Sbp1* under normal conditions and in response to sodium azide stress.

Wild type cells showed modulation of many transcripts in response to sodium azide treatment. Thirteen genes were significantly

upregulated and belonged to the GO biological process category of ‘response to oxidative stress’ (Fig. 4a and Supplementary Fig. 4b & c). In this category, genes involved in protection against oxidative damage (CTT1, SRX1, CCP1, GAD1, and NCE103) and multi-stress response (HSP30, MTL11, and DDR2) were strongly induced. These results

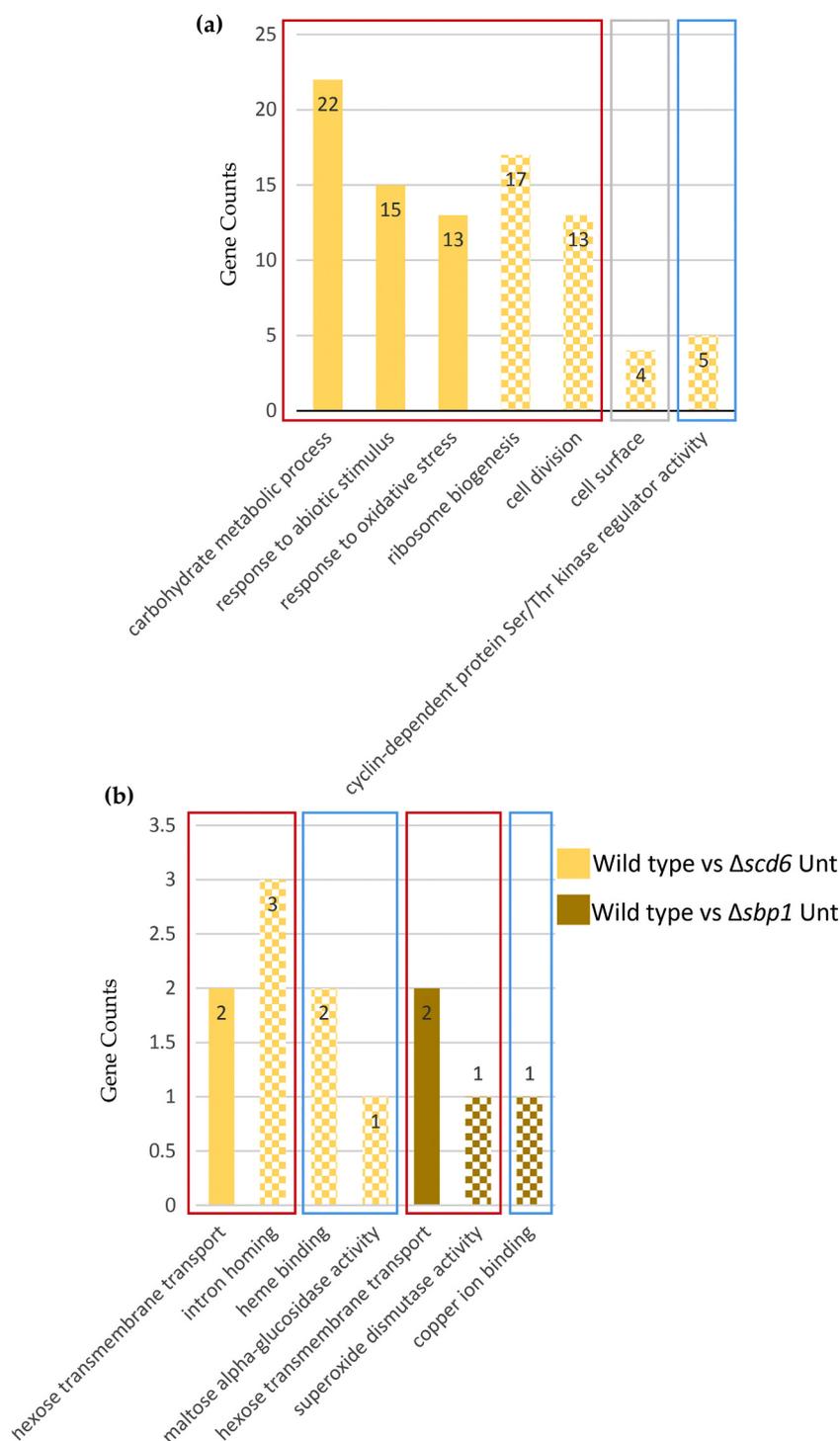


Fig. 4. Gene ontology enrichment analysis of differentially expressed genes in (a) wild type untreated vs treated cells and (b) wild type vs $\Delta scd6/\Delta sbp1$ untreated cells. Solid and pattern fill indicate up and downregulated categories, respectively. Red, blue and grey rectangular boxes represent Biological Processes, Molecular Function and Cellular Component, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

emphasized the role of sodium azide as an agent of oxidative damage to the cells. Cytosolic Catalase T (CTT1) is known for its role in the oxidative stress response. Ctt1 activity is crucial to protect proteins from oxidative damage [42] and CTT1 upregulation is reported in response to a number of stresses such as starvation, osmotic, and oxidative stress [43]. A similar increase in CTT1 mRNA level was observed in our study, highlighting a feature of stress response mounted by yeast cells to counter oxidative damage induced by sodium azide.

Transcripts encoding proteins associated with carbohydrate metabolism were induced in stressed cells (Fig. 4a and Supplementary Fig. 4b & c). This response could fulfil the energy requirements during the stress response. Some of these upregulated genes belong to the glycolytic pathway (TDH1, GPM2, and ENO1) while others were related to closely associated pathways (YIG1, GCY1, GID8, and GPH1). Glycogen and trehalose are the two storage forms of glucose. Our transcriptome analysis also showed upregulation of transcripts (GLC3,

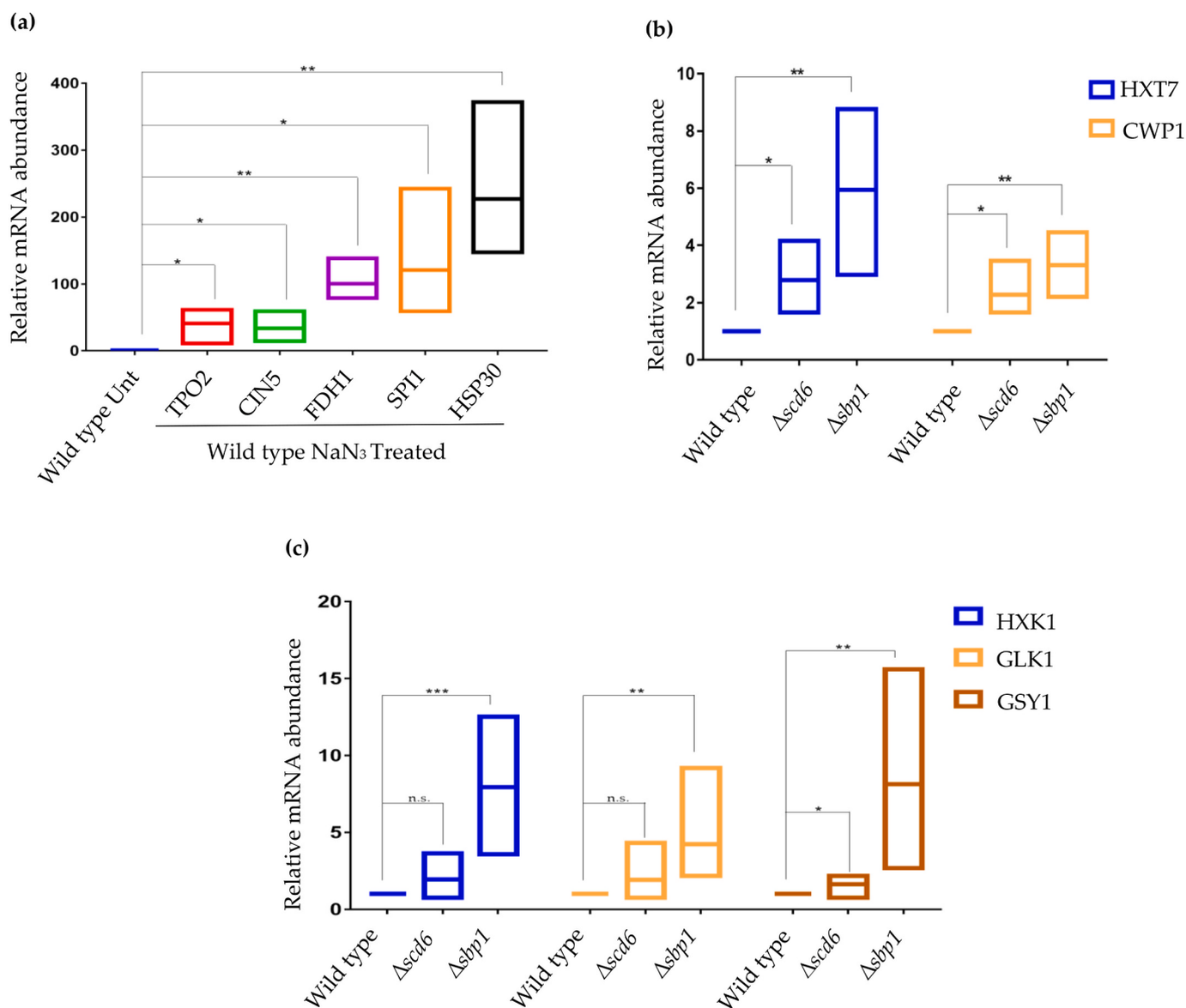


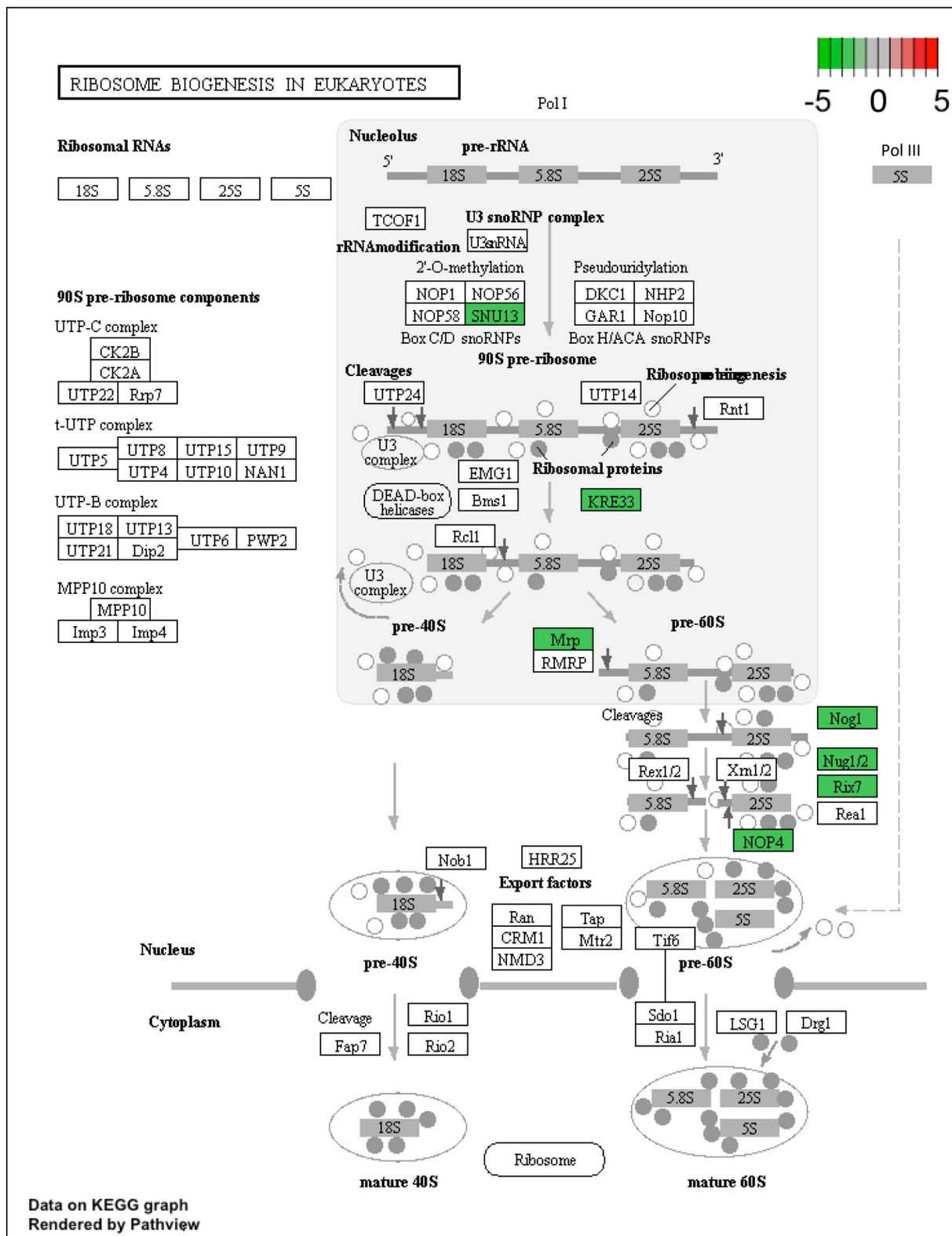
Fig. 5. Validation of differentially expressed genes by qRT-PCR analysis in (a) wild type untreated vs NaN_3 treated cells and (b) wild type vs $\Delta\text{scd6}/\Delta\text{sbp1}$ cells and (c) wild type vs $\Delta\text{scd6}/\Delta\text{sbp1}$ cells. Internal control for (a) and (b) and (c) are PGK1 and SCR1, respectively ($n > 3$; *, ** and *** denotes p -value $< .05$, p -value $< .01$ and p -value $< .001$ respectively).

IGD1, GSY2, and TSL1) encoding enzymes catalyzing the synthesis of these two polysaccharides. This is in agreement with the reported observation that glycogen and trehalose production are elevated in other stresses [44].

Genes involved in gamma-aminobutyric (GABA) acid metabolism (belonging to the Butanoate pathway) play a role in the oxidative stress response to stressors such as H_2O_2 and diamide [45]. Diamide depletes glutathione and oxidize thiol groups. Two genes of this pathway, GAD1 and UGA2, were upregulated upon sodium azide treatment in wild type, Δscd6 and Δsbp1 cells, indicating their role in the sodium azide stress response. The pentose phosphate pathway (PPP) is also upregulated in response to oxidative stress [46]. SOL4, PGM2, and NQM1 are PPP components upregulated in wild type cells in response to sodium azide stress in our study. This increase would generate NADPH to provide reducing potential and act as a cofactor for glutathione- and thiorodoxin-dependent enzymes that play a major role in survival during the oxidative stress response [47].

Downregulation of cellular processes that consume a high amount of energy, such as translation and cell cycle progression, is a feature of

stress response pathways. We find evidence supporting this in the sodium azide stress response program. In our study, multiple genes in the ribosome biogenesis pathway (SNU13, KRE33, MRP, NOG1, NUG1/2, RIX7, and NOP4) were downregulated following sodium azide treatment (Fig. 6). Ribosome biogenesis is a highly energy-expensive process and defects in it lead to translation repression [48]. Our observation is highly consistent with the report that sodium azide treatment leads to global translation repression and the assembly of cytoplasmic repression foci [49]. Similarly, cell cycle genes such as PCL1, CLN1/2, CLB5/6, CDC6, HSL1 were downregulated upon azide treatment (Supplementary Fig. 8a). This is in accordance with the observation that oxidative stress downregulates expression of cell cycle genes involved in the G1 to S transition [50]. Together, the above observations indicate that the response to sodium azide stress is similar to other oxidative stress response pathways. Interestingly, though ribosome biogenesis related genes were downregulated in wild type, Δscd6 and Δsbp1 background upon sodium azide treatment, the cell cycle related target genes were downregulated in wild type and Δsbp1 background but not in Δscd6 cells (Supplementary Fig. 4b & <https://www.ncbi.nlm.nih>).



Ribosome biogenesis pathway

Fig. 6. Ribosome biogenesis pathway showing genes which are significantly changing upon sodium azide treatment of wild type cells. The scale is set as log2fold change where downregulated genes are shaded green. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE145162>.

Some GO categories were found to be enriched only in the case of deletion of Scd6 and Sbp1 upon sodium azide treatment

(Supplementary Fig. 4b & c). This reflects the importance of these proteins in mounting an appropriate cell response to stress.

The cell wall is a major barrier to physical stressors in yeast [51].

Cell wall synthesis and repair enzymes are activated in response to oxidative stress such as H₂O₂. We capture this behavior in our study in response to sodium azide treatment. Genes involved in cell wall integrity pathways, such as SDPI, MID2, and MTL1, are upregulated upon sodium azide treatment, as has been observed with other oxidative stressors [52].

Many mitogen-activated protein kinase (MAPK) pathway genes are downregulated in response to sodium azide stress, including MSB2, HSL1, BAR1, SPA2, YPS1, and YPS2. This is surprising given that the MAPK pathway is activated upon H₂O₂-mediated oxidative stress [53]. This observation indicates that although shared metabolic regulators are elicited by sodium azide and H₂O₂, the signaling pathways activated in response to these two stressors may be distinct.

Downregulation of purine and pyrimidine biosynthesis pathway genes (HPRT1, RNRI, and URA2) likely indicate a stress response mechanism aimed at reducing nucleotide biosynthesis during oxidative stress (Supplementary Fig. 9).

We validated the modulation in the expression levels of HSP30, SPI1, TPO2, CIN5, and FDH1 genes in wild type cells subjected to sodium azide treatment (Fig. 5a). HSP30 and SPI1 were previously reported to be stress-responsive genes, albeit not in response to sodium azide stress. HSP30 is induced by heat shock, ethanol treatment, and weak organic acid. SPI1 is a GPI-anchored cell wall protein (encoded by SPI1 mRNA) induced by stress conditions and contributes to weak acid resistance [54]. FDH1 encodes formate dehydrogenase, and is induced by the presence of formate in the media [55]. TPO2 encodes a spermine-specific polyamine transporter that localizes to the plasma membrane [56]. All of the above genes were not reported to change in response to oxidative stress such as H₂O₂, indicating that the stress response mediated through these genes may not overlap with the H₂O₂ stress response, thereby supporting the idea that cells use different sets of proteins to respond to different stressors. In our study, upregulation of these transcripts upon sodium azide treatment indicates their involvement in the sodium azide stress response. The specific contribution of these genes to the sodium azide stress response will be evaluated in future.

RNA binding proteins play a critical role in regulating mRNA fate and abundance. Our report studied the importance of two RNA binding proteins, Scd6 and Sbp1, which are known translation repressors, decapping activators, and RNA granule modulators [23,25,26,57]. Three meiosis/sporulation specific genes were downregulated in $\Delta scd6$ including SPO19, SPO23, and AMA1 (Supplementary Table 10). Spo19 and Spo23 are pro-spore and Spo1-associated protein, respectively [58,59]. Ama1 is an activator of the meiotic anaphase-promoting complex, which is required to initiate spore wall assembly during anaphase [60,61]. These observations suggest that Scd6 could be involved in sporulation and/or meiosis related processes, which has not been addressed previously in the literature. DAN1 transcript expression increases in $\Delta scd6$ cells. DAN1 is a cell wall mannoprotein expressed under anaerobic conditions and repressed under aerobic conditions [62]. Although the precise role of DAN1 in anaerobic growth is not clear, it is interesting to note that the yeast strain lacking Scd6 is sensitive to oxygen deprivation stress [63]. The mechanisms underlying changes in DAN1 expression level and $\Delta scd6$ sensitivity to anaerobic stress is an interesting aspect to study in future. Strikingly, many transcripts encoding proteins of unknown function (PUF) were differentially expressed in the absence of Scd6 (refer to the deposited data, GEO: GSE145162). The functional significance of these changes is currently unclear. Studying the function of Scd6 in detail could reveal the function of genes encoding PUFs. Overall, the transcripts upregulated in $\Delta scd6$ formed two clusters (Supplementary Fig. 11). The first cluster contains eight genes with known or putative helicase functions, including YRF1–5 and YRF1–4. The second cluster includes transporter proteins, such as sugar transporter genes like HXT6 and HXT7, and sodium pump ENA2.

The decapping activator Sbp1 binds mRNA, likely in the 5'UTR

region [30]. Sbp1 deletion upregulated some genes, including HXK1 and HXT7 (Supplementary Fig. 6c), which are normally downregulated in the presence of glucose. Other genes, such as GSY1, GSY2, ALD4, and GLK1 that are downregulated in the presence of glucose were also found to be upregulated when the threshold value of log₂FC was lowered to > 1 from > 2. HXK1 and GLK1 transcripts both encode for kinases that phosphorylate glucose in the first step of glycolysis. Expression of both the genes is repressed in the presence of glucose and de-repressed in the presence of non-fermentable carbon sources and galactose [64]. HXK2, a paralog of HXK1, is a predominant hexokinase that promotes repression of HXK1 and GLK1 expression in glucose containing glucose [65]. It was recently reported that Sbp1 overexpression from a galactose-inducible promoter leads to a growth defect on galactose media [25]. The growth defect could be due to sustained translation repression of HXK1 and GLK1 by Sbp1 overexpression, leading to less availability of non-glucose specific kinases. It would be important to test if the lack of de-repression of HXK1 and GLK1 upon Sbp1 overexpression in galactose media contributes to the growth defect phenotype.

A feature of a subset of differentially regulated mRNAs in $\Delta sbp1$ was their association with mitochondria. This included transcripts encoding mitochondrial enzymes like aldehyde dehydrogenase (ALD4) and glyoxylate reductase (GOR1), and mitochondrial membrane proteins like FMP45 and CYC7. The connection between Sbp1 and mitochondria-related transcripts would be an important direction to pursue in future studies.

Genes upregulated in $\Delta sbp1$ include stress responsive genes such as DDR2 and HSP12. Interestingly, these genes were also upregulated in wild type cells in response to sodium azide treatment. A similar observation was made with carbohydrate metabolic process genes such as GSY2, GPM2, GPH1, and SOL4. Based on the above observations, it is likely that these genes are repressed by Sbp1 under normal conditions and derepressed in response to sodium azide stress. This de-repression could be orchestrated by post-translational modifications of Sbp1, such as arginine methylation. Interestingly, Sbp1 methylation, which is increased upon glucose deprivation, promotes translation repression and decapping [25].

Comparing the sodium azide stress response in wild type and $\Delta scd6$ or wild type and $\Delta sbp1$ strains revealed differential expression in several classes of transcripts (Supplementary Fig. 4). In $\Delta scd6$ as compared to wild type in azide treatment, transcripts encoding cell cycle category proteins were significantly upregulated (Supplementary Fig. 8b). For example, PCL1/2, CLN1/2, and CLB5/6, which were otherwise downregulated in the wild type untreated vs treated condition (Supplementary Fig. 8a). Other interesting observation which came out was the differential regulation of MAL32, an inducible protein involved in maltase catabolism. In $\Delta scd6$ the abundance of MAL32 transcript reduced significantly as compared to in wild type (Supplementary Fig. 10) but upon treatment of $\Delta scd6$ cells with sodium azide, transcript levels got enriched. These observations suggest that Scd6 might play a role in controlling the abundance of different transcripts during azide stress in a different manner. In the case of $\Delta sbp1$ sodium azide treated cells as compared to wild type treated cells, one of the subunits of cytochrome C oxidase (COX1) belonging to the GO category of aerobic electron transport chain (Supplementary Fig. 4c) is significantly upregulated (Supplementary Fig. 10). This is interesting given that COX1 mRNA was downregulated in wild type cells following sodium azide treatment (Supplementary Fig. 10). Further, Cox1 is a mitochondrially-encoded protein. Understanding the mechanism underlying differential modulation of mitochondrial transcripts by cytosolic translation repressor proteins will be an important future direction.

Comparative analysis of transcripts differentially expressed in response to sodium azide stress in wild type, $\Delta scd6$, and $\Delta sbp1$ strains (Fig. 7) indicated that 120 transcripts change in abundance only in $\Delta sbp1$. Similarly, 204 transcripts changed in abundance in $\Delta scd6$, but not in wild type or $\Delta sbp1$. Based on this, we propose that the absence of

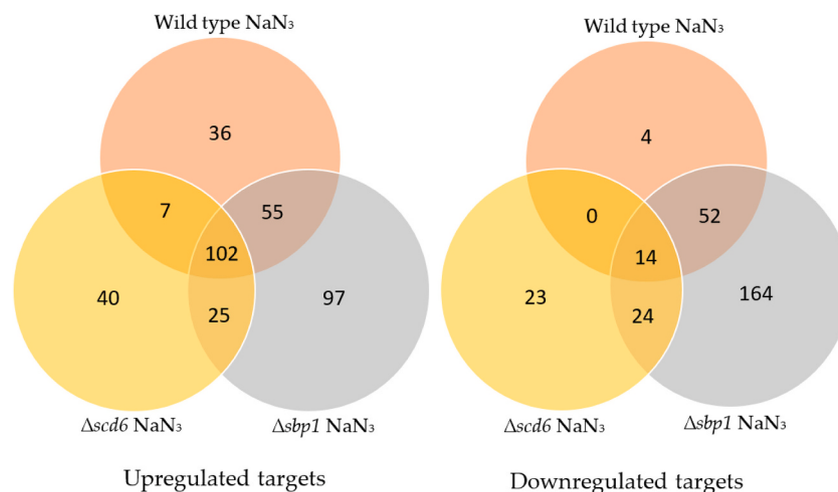


Fig. 7. Venn diagrams showing number of genes that are differentially expressed upon sodium azide treatment in wild type, $\Delta scd6$ and $\Delta sbp1$ cells.

Sbp1 or Scd6 affects the transcriptome differentially in response to sodium azide stress. We observe that the number of genes differentially changing between wild type and $\Delta sbp1$ cells upon sodium azide treatment is much smaller (Fig. 2) than the wild type (untreated vs treated) and $\Delta sbp1$ (untreated vs treated). We interpret this to indicate that global expression level changes are similar in wild type and $\Delta sbp1$ backgrounds when azide treatment was given. This also seems to be the case between wild type and $\Delta scd6$ background. Our study provides a critical starting point to address the mechanistic details of the sodium azide stress response in yeast. The information provided by this study will allow hypothesis-driven dissection of the sodium azide stress response in general and specifically regarding translation repression and RNA granule assembly. The observations presented here identify the mRNA targets of Scd6 and Sbp1. Interestingly out of all the genes validated, two genes, HXT7 and CWP1, are affected by deletion of both Scd6 and Sbp1, although to different extents (Fig. 5b and Supplementary Fig. 6b & c). It is possible that both proteins act in conjunction to affect these mRNAs. It will be interesting to investigate the effect of $\Delta sbp1\Delta scd6$ double deletion on the commonly affected genes. As discussed in the introduction, since both the proteins can act as decapping activators, it is likely that the upregulated transcripts are a result of increased mRNA stability. Identifying mRNA decay targets of Scd6 and Sbp1 from the upregulated transcripts will be an obvious future endeavor. It is intriguing that the number of downregulated transcripts in the absence of these decapping activators are comparable to the number of upregulated transcripts (Fig. 2). Scd6 orthologs play a role in protection and maintenance of certain developmental mRNAs in *Drosophila* and *C. elegans* [24,32,33]. Upon depletion of Scd6 orthologs, the repression complex is hampered, leading to the loss of mRNA-protection in germ granules, making those transcripts susceptible to degradation. In our study, the genes downregulated in $\Delta sbp1$ and $\Delta scd6$ could be due to a similar loss of mRNA masking mechanisms. Such mRNAs could become substrates for mRNA degradation and hence get downregulated. It is possible that Scd6/Sbp1 could also affect transcription of some of these mRNAs either directly or indirectly. mRNA degradation can affect transcription, and some decay factors have been implicated in directly modulating transcription [66]. Interestingly, Sbp1 (earlier referred to as Ssb1) was reported as a nucleolar protein that can bind snRNAs based on studies using anti-Sbp1 antibodies [67]. However, GFP-tagged Sbp1 is uniformly localized in yeast cells [25]. Whether this is due to differences pertaining to live cell imaging versus fixed cell staining or due to GFP-tagging remains to be addressed. Many RGG-motif proteins, such as Npl3 and Gbp2, are nucleo-cytoplasmic shuttling proteins and are known to affect nuclear processes [23]. The ability of Scd6/Sbp1 to shuttle in and out of the nucleus followed by its

impact on the target mRNAs identified in this study will be addressed in our future studies.

The following is the supplementary data related to this article.

The data discussed in this publication has been deposited in NCBI's Gene Expression Omnibus [68] and are accessible through GEO accession number: [GSE145162](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE145162) (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE145162>).

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Declaration of Competing Interest

The authors declare no conflict of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.ygeno.2020.05.001>.

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